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THOMAS CAV	7590 01/16/2007 VLEY		EXAM	INER
Pillsbury Winthrop, LLP			CORDERO GARCIA, MARCELA M	
1600 Tysons Blvd. McLean, VA 22102		· ·	ART UNIT	PAPER NUMBER
,			1654	
SHORTENED STATUTORY PERIOD OF RESPONSE		MAIL DATE	DELIVED	Y MODE
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Please find below and/or attached an Office communication concerning this application or proceeding.

If NO period for reply is specified above, the maximum statutory period will apply and will expire 6 MONTHS from the mailing date of this communication.

	Application No.	Applicant(s)				
Office Action Comment	10/796,158	BRASLAWSKY ET AL.				
Office Action Summary	Examiner	Art Unit				
÷	Marcela M. Cordero Garcia	1654				
The MAILING DATE of this communication appears on the cover sheet with the correspondence address Period for Reply						
A SHORTENED STATUTORY PERIOD FOR REPLY IS SET TO EXPIRE 3 MONTH(S) OR THIRTY (30) DAYS, WHICHEVER IS LONGER, FROM THE MAILING DATE OF THIS COMMUNICATION. - Extensions of time may be available under the provisions of 37 CFR 1.136(a). In no event, however, may a reply be timely filed after SIX (6) MONTHS from the mailing date of this communication. - If NO period for reply is specified above, the maximum statutory period will apply and will expire SIX (6) MONTHS from the mailing date of this communication. - Failure to reply within the set or extended period for reply will, by statute, cause the application to become ABANDONED (35 U.S.C. § 133). Any reply received by the Office later than three months after the mailing date of this communication, even if timely filed, may reduce any earned patent term adjustment. See 37 CFR 1.704(b).						
Status						
1) Responsive to communication(s) filed on 20 Oc	Responsive to communication(s) filed on 20 October 2006.					
· · · · · · · · · · · · · · · · · · ·						
·= ·	Since this application is in condition for allowance except for formal matters, prosecution as to the merits is					
closed in accordance with the practice under E	closed in accordance with the practice under Ex parte Quayle, 1935 C.D. 11, 453 O.G. 213.					
Disposition of Claims						
4)⊠ Claim(s) <u>1-20</u> is/are pending in the application.						
4a) Of the above claim(s) <u>1-14</u> is/are withdrawn from consideration.						
5) Claim(s) is/are allowed.						
6)⊠ Claim(s) <u>15-20</u> is/are rejected.						
7) Claim(s) is/are objected to.		·				
8) Claim(s) are subject to restriction and/or	election requirement.					
Application Papers						
9)☐ The specification is objected to by the Examiner.						
10)☐ The drawing(s) filed on is/are: a)☐ accepted or b)☐ objected to by the Examiner.						
Applicant may not request that any objection to the drawing(s) be held in abeyance. See 37 CFR 1.85(a).						
Replacement drawing sheet(s) including the correction is required if the drawing(s) is objected to. See 37 CFR 1.121(d).						
11) The oath or declaration is objected to by the Examiner. Note the attached Office Action or form PTO-152.						
Priority under 35 U.S.C. § 119						
12) Acknowledgment is made of a claim for foreign priority under 35 U.S.C. § 119(a)-(d) or (f). a) All b) Some * c) None of:						
1. Certified copies of the priority documents have been received.						
2. Certified copies of the priority documents have been received in Application No						
3. Copies of the certified copies of the priority documents have been received in this National Stage						
application from the International Bureau (PCT Rule 17.2(a)).						
* See the attached detailed Office action for a list of the certified copies not received.						
		·				
Attachment(s)						
1) Notice of References Cited (PTO-892)	4) Interview Summary	(PTO-413)				
2) Notice of Draftsperson's Patent Drawing Review (PTO-948)	Paper No(s)/Mail Da	te				
3) Information Disclosure Statement(s) (PTO/SB/08) Paper No(s)/Mail Date 5) Notice of Informal Patent Application 6) Other:						
	-/ <u></u>					

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DETAILED ACTION

This Office Action is in response to the reply received on October 20, 2005.

Claims 1-20 are pending in the application. Claims 1-14 have been withdrawn.

Any rejection from the previous office action, which is not restated here, is withdrawn.

Claim 15 has been amended.

Applicants elected without traverse claims 15-20 (Group III), directed to methods of treating an SSTR-associated disorder. In response to the election of species requirement, Applicants elected the somatostatin analog wherein A is SEQ ID NO: 1 (claim 16) and B is SEQ ID NO:4 (claim 17), which is equivalent to the CP1 somatostatin analog of SEQ ID NO: 5 (claim 18).

The species was searched and found free of the prior art. However, please note that no claims are drawn uniquely to this single species. The search was broaden to encompass claims SEQ IDs NOs: 1-7, which were found free of the prior art. The search was extended by Examiner to the species:

wherein A is cysteine or a peptide chain comprising one or more cysteine residues; R=H, wherein paclitaxel (a therapeutic agent) is bound to the cysteine residue via a thiol linkage and B= octreotide (D-Phe-Cys-Phe-DTrp-Lys-Thr-Cys-Thr(ol). The previous 103 rejection over this species has been overcome by Applicant's amendments to claim 15.

The search has been broadened again and claims 15-20 are presented for examination on the merits.

Claim Rejections - 35 USC § 112

The following is a quotation of the first paragraph of 35 U.S.C. 112:

The specification shall contain a written description of the invention, and of the manner and process of making and using it, in such full, clear, concise, and exact terms as to enable any person skilled in the art to which it pertains, or with which it is most nearly connected, to make and use the same and shall set forth the best mode contemplated by the inventor of carrying out his invention.

Claims 15-20 are rejected under 35 U.S.C. 112, first paragraph, as failing to comply with the written description requirement. The claim(s) contains subject matter which was not described in the specification in such a way as to reasonably convey to one skilled in the relevant art that the inventor(s), at the time the application was filed, had possession of the claimed invention. The MPEP states that the purpose of the written description requirement is to ensure that the inventor had possession, as of the filing date of the application, of the specific subject matter later claimed by him. The courts have stated:

"To fulfill the written description requirement, a patent specification must describe an invention and do so in sufficient detail that one skilled in the art can

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clearly conclude that "the inventor invented the claimed invention." Lockwood v. American Airlines, Inc., 107 F.3d 1565, 1572, 41 USPQ2d 1961, 1966 (1997); In re Gosteli, 872 F.2d 1008, 1012, 10 USPQ2d 1614, 1618 (Fed. Cir. 1989) (" [T]he description must clearly allow persons of ordinary skill in the art to recognize that [the inventor] invented what is claimed."). Thus, an applicant complies with the written description requirement "by describing the invention, with all its claimed limitations, not that which makes it obvious," and by using "such descriptive means as words, structures, figures, diagrams, formulas, etc., that set forth the claimed invention." Lockwood, 107 F.3d at 1572, 41 USPQ2d at 1966." Regents of the University of California v. Eli Lilly & Co., 43 USPQ2d 1398.

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The MPEP lists factors that can be used to determine if sufficient evidence of possession has been furnished in the disclosure of the Application. These include "level of skill and knowledge in the art, partial structure, physical and/or chemical properties, functional characteristics alone or coupled with a known or disclosed correlation between structure and function, and the method of making the claimed invention. Disclosure of any combination of such identifying characteristics that distinguish the claimed invention from other materials and would lead one of skill in the art to the conclusion that the applicant was in possession of the claimed species is sufficient." MPEP 2163.

Further, for a broad generic claim, the specification must provide adequate written description to identify the genus of the claim. In Regents of the University of California v. Eli Lilly & Co., the court stated:

"A written description of an invention involving a chemical genus, like a description of a chemical species, 'requires a precise definition, such as by structure, formula, [or] chemical name,' of the claimed subject matter sufficient to distinguish it from other materials.

5 USPQ2d at 1606; In re Smythe, 480 F.2d 1376, 1383, 178 USPQ 279, 284-85 (CCPA 1973) ("In other cases, particularly but not necessarily, chemical cases, where there is unpredictability in performance of certain species or subcombinations other than those specifically enumerated, one skilled in

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the art may be found not to have been placed in possession of a genus. . . ."). Regents of the University of California v. Eli Lilly & Co., 43 USPQ2d 1398.

The MPEP further states that if a biomolecule is described only by a functional characteristic, without any disclosed correlation between function and structure of the sequence (e.g., "a therapeutic agent"), it is "not sufficient characteristic for written description purposes, even when accompanied by a method of obtaining the claimed sequence." MPEP 2163. The MPEP does state that for generic claim the genus can be adequately described if the disclosure presents a sufficient number of representative species that encompass the genus. MPEP 2163. If the genus has a substantial variance, the disclosure must describe a sufficient variety of species to reflect the variation within that genus. See MPEP 2163. Although the MPEP does not define what constitute a sufficient number of representative, the Courts have indicated what do not constitute a representative number species to adequately describe a broad generic. In Gostelli, the Court determined that the disclosure of two chemical compounds within a subgenus did not describe that subgenus. In re Gostelli, 872 F.2d at 1012, 10 USPQ2d at 1618.

In the instant case, the claims are drawn to a method of treating an SSTR-associated disorder in a mammalian subject comprising administering to the subject a composition comprising (A-B) wherein A is a peptide chain comprising one or more cysteine residues and B is a naturally occurring or synthetic somatostatin peptide or fragment thereof, which binds to a somatostatin receptor, wherein a therapeutic agent is bound to the somatostatin analog (A-B) via a thiol linkage to the one or more cysteine residues of (A) at an interior site(s); whereby a SSTR is treated. In regards to the "somatostatin peptide or fragment thereof" term, this is a very broad generic definition, which is not adequately described

and/or represented in the examples. Please note that the specification provides only one example, with a generic CP1 as somatostatin analog, for which the formula is not described, with a derivative of aurestatin E. By the same token, the peptides comprise the instant SEQ ID Nos: 1-7, however, no guidance is provided in terms of the sequence related to such larger peptides encompassing SEQ ID Nos: 1-7 to be used in the instant methods, nor does it provide teachings for how to find the variety of peptide derivatives, mutations, variants, analogs, fragments, peptoids, chemically modified peptides thereof as mentioned in the instant disclosure, page 9. In addition therapeutic agents (e.g., disclosure, pages 18-20) include any agents with activity against cancer or SSTR-related disease, therapeutic genes, immunostimulatory agents and so forth, therefore a mere statement that such compounds would be desirable for conjugation does not sufficiently provide ample written description pages describing the full breadth of the (A-B) thiol containing conjugates for the biological activity of the instantly claimed method. The specification does provide examples of what qualify as compounds of the claimed invention (see, e.g., disclosure, pages), however, these are limited to a few examples such CP1-AEB, CP1-FKMMAE and CP1-MX-DTPA (e.g., pages 27-30) Please note that pages 18-20 describe various desirable therapeutic agents but there is no conjugation therein with the instantly claimed compounds of formula (A-B) and only one example wherein the biological activity of the instantly claimed conjugates is tested. As stated earlier, the MPEP states that written description for a genus can be achieved by a representative number of species within a broad generic. It is unquestionable claim 15 is a broad generic with respect all possible compounds encompassed by the claims. The possible structural variations are very broad as the compound would comprise A-B but there is no restriction in regards to how A binds B and/or the many possible somatostatin analog prodrugs comprising this

generic structure and elements. It must not be forgotten that the MPEP states that if a biomolecule, if described only by a functional characteristic, without any disclosed correlation between function and structure of the sequence, it is "not sufficient characteristic for written description purposes, even when accompanied by a method of obtaining the claimed sequence." MPEP 2163. Here, though the claims recite some functional and structural characteristics, the claims lack written description because there is no disclosure of a correlation between function and structure of the compounds beyond compounds disclosed in the examples in the specification. Moreover, the specification lack sufficient variety of species to reflect this variance in the genus since the specification does not provide any examples of conjugates with e.g., chemically modified peptides, peptoids, mutations, variants analogs, peptide derivatives, with therapeutic agents such as antiangiogenic agents, therapeutic genes, immunostimultory agents, anticancer agents and so forth. The description requirement of the patent statute requires a description of an invention, not an indication of a result that one might achieve if one made that invention. See In re Wilder, 736 F.2d 1516, 1521, 222 USPQ 369, 372-73 (Fed. Cir. 1984) (affirming rejection because the specification does "little more than outlin[e] goals appellants hope the claimed invention achieves and the problems the invention will hopefully ameliorate."). Accordingly, it is deemed that the specification fails to provide adequate written description for the genus of the claims and does not reasonably convey to one skilled in the relevant art that the inventor(s), at the time the application was filed, had possession of the entire scope of the claimed invention.

The following is a quotation of the second paragraph of 35 U.S.C. 112:

The specification shall conclude with one or more claims particularly pointing out and distinctly claiming the subject matter which the applicant regards as his invention.

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Claims 15-16 are rejected under 35 U.S.C. 112, second paragraph, as being indefinite for failing to particularly point out and distinctly claim the subject matter which applicant regards as the invention.

Claim 16 is vague and indefinite by the phrase in line 2: "wherein A is a single cysteine residue" because claim 15, of which it depends, defines "A is a peptide chain" which necessitates at least a few amino acid residues and not just a single cysteine residue.

Claim Rejections - 35 USC § 102

The following is a quotation of the appropriate paragraphs of 35 U.S.C. 102 that form the basis for the rejections under this section made in this Office action:

A person shall be entitled to a patent unless -

(b) the invention was patented or described in a printed publication in this or a foreign country or in public use or on sale in this country, more than one year prior to the date of application for patent in the United States.

Claim 15 is rejected under 35 U.S.C. 102(b) as being anticipated by Fukuta et al. (US 5,442,043).

A method for treating an SSTR-associated disorder in a mammalian subject (column 3, lines 13-56, column 8, lines 10-45), the method comprising administering to the subject a composition (e.g., column 14, lines 17-24) comprising a somatostatin analog of the formula (A-B) wherein A is a peptide chain comprising one or more cysteine residues (e.g., A is F007 as in Figure 6 and Example 10; column 13, lines 36-40; column 8, lines 10-45) and B is somatostatin (e.g., Example 10).

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Please note that according to MPEP 2111.02, the determination of whether a preamble limits a claim is made on a case-by-case basis in light of the facts in each case; there is no litmus test defining when a preamble limits the scope of a claim.

Catalina Mktg. Int'l v. Coolsavings.com, Inc., 289 F.3d 801, 808, 62 USPQ2d 1781, 1785 (Fed. Cir. 2002). See id. at 808-10, 62 USPQ2d at 1784-86 for a discussion of guideposts that have emerged from various decisions exploring the preamble's effect on claim scope, as well as a hypothetical example illustrating these principles.

"[A] claim preamble has the import that the claim as a whole suggests for it." *Bell Communications Research, Inc. v. Vitalink Communications Corp.*, 55 F.3d 615, 620, 34 USPQ2d 1816, 1820 (Fed. Cir. 1995). "If the claim preamble, when read in the context of the entire claim, recites limitations of the claim, or, if the claim preamble is 'necessary to give life, meaning, and vitality' to the claim, then the claim preamble should be construed as if in the balance of the claim." *Pitney Bowes, Inc. v. Hewlett-Packard Co.*, 182 F.3d 1298, 1305, 51 USPQ2d 1161, 1165-66 (Fed. Cir. 1999). See also *Jansen v. Rexall Sundown, Inc.*, 342 F.3d 1329, 1333, 68 USPQ2d 1154, 1158 (Fed. Cir. 2003)(In considering the effect of the preamble in a claim directed to a method of treating or preventing pernicious anemia in humans by administering a certain vitamin preparation to "a human in need thereof," the court held that the claims' recitation of a patient or a human "in need" gives life and meaning to the preamble's statement of purpose.).

In regards to the "wherein" and "whereby" clauses (MPEP 2111.04) claim scope is not limited by claim language that suggests or makes optional but does not require steps to be performed, or by claim language that does not limit a claim to a particular structure. However, examples of claim language, although not exhaustive, that may raise a question as to the limiting effect of the language are "wherein" and "whereby".

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The determination of whether each of these clauses is a limitation in a claim depends on the specific facts of the case. In *Hoffer v. Microsoft Corp.*, 405 F.3d 1326, 1329, 74 USPQ2d 1481, 1483 (Fed. Cir. 2005), the court held that when a "whereby' clause states a condition that is material to patentability, it cannot be ignored in order to change the substance of the invention." *Id.* However, the court noted (quoting *Minton v. Nat'l Ass'n of Securities Dealers, Inc.*, 336 F.3d 1373, 1381, 67 USPQ2d 1614, 1620 (Fed. Cir. 2003)) that a "whereby clause in a method claim is not given weight when it simply expresses the intended result of a process step positively recited." *Id.*<

In the instant claims, the conjugate insulin-somatostatin can by itself treat a SSTR disease, and therefore the preamble and the whereby clauses are not deemed to modify the claims structurally. The wherein clause is also not deemed to be limiting, please note that Fig. 6 shows the insulin has thiol linkages between the insulin A and B chains, and that each one can be deemed a therapeutic agent (see also claims 1 and 10 of Fukuta et al.).

Therefore, the reference is deemed to anticipate the instant claims above.

Conclusion

No claim is allowed.

The prior art made of record and not relied upon is considered pertinent to applicant's disclosure.

Any inquiry concerning this communication or earlier communications from the examiner should be directed to Marcela M. Cordero Garcia whose telephone number is (571) 272-2939. The examiner can normally be reached on M-Th 7:30-6:00.

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If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, Cecilia J. Tsang can be reached on (571) 272-0562. The fax phone number for the organization where this application or proceeding is assigned is 571-273-8300.

Information regarding the status of an application may be obtained from the Patent Application Information Retrieval (PAIR) system. Status information for published applications may be obtained from either Private PAIR or Public PAIR. Status information for unpublished applications is available through Private PAIR only. For more information about the PAIR system, see http://pair-direct.uspto.gov. Should you have questions on access to the Private PAIR system, contact the Electronic Business Center (EBC) at 866-217-9197 (toll-free). If you would like assistance from a USPTO Customer Service Representative or access to the automated information system, call 800-786-9199 (IN USA OR CANADA) or 571-272-1000.

Marcela M Cordero Garcia, PhD

aM Corden Jano

Patent Examiner

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